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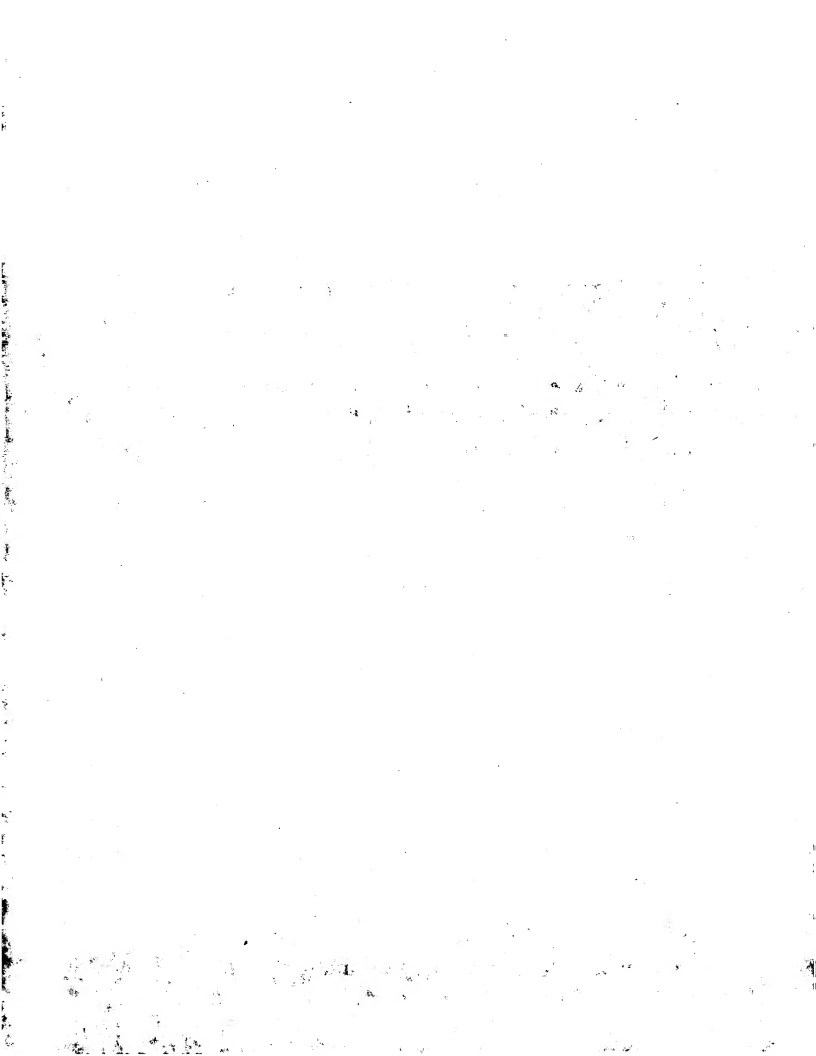
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PATENT SPECIFICATION



NO DRAWINGS

1011976

Date of Application and filing Complete Specification: May 10, 1962. No. 18060/62.

Application made in United States of America (No. 109,007) on May 10, 1961. Application made in United States of America (No. 186,325) on April 10, 1962. Complete Specification Published: Dec. 1, 1965.

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Index at acceptance:—C3 R(3C8, 3C9, 3C12, 3C16, 3C17, 3D2A, 3D2B, 3D3, 3D5, 3D10, 3D11, 3D17, 3D18, 29C8, 29C9, 29C12, 29C17, 33C9, 33C12, 33P); C2 C(1Q2, 1Q6A1, 1Q8A, 1Q9A, 1Q9D1, 1Q9D2, 1Q11G, 1Q11J); C3 B(1C8, 1C9, 1C12, 1C17, 1D1X, 1D2C, 1N1F, 1N2A, 1N4B, 1N4C, 1N4F, 1N4H, 1N4J, 1N6D, 1N6X, 1N7, 1N9X, 1N10, 1N15, 1N16B); C3 P(14D1A, 14D1B, 14D2B, 14D2F, 14D2G1, 14D2G2, 14D2H, 14D3A2, 14D3C1, 14D4, 14D5, 14D6, 14D7X, 14K4, 14K7, 14K8, 14P1B, 14P1E1)

Int. Cl.:—C 08 g // C 07 c, C 08 f

COMPLETE SPECIFICATION

Fluorine-Containing Resins

We, MINNESOTA MINING AND MANUFAC-TURING COMPANY, a Corporation organised and existing under the laws of the State of Delaware, United States of America, of 900 Bush Avenue, Saint Paul 6, Minnesota, United States of America, do hereby declare the invention, for which we pray that a patent may be granted to us and the control of the c patent may be granted to us, and the method by which it is to be performed, to be par-10 ticularly described, in and by the following statement:-

This invention relates to new and useful fluorine-containing resins and their method of preparation. In one aspect this invention 15 relates to modified resin compositions and their use in varnishes, lacquers, paints and various coating compositions.

Fluorocarbon compounds have been known to impart water and oil repellency to various materials and have been used to treat such materials as fabrics, leather, wood, kapok, paper, fur, asbestos, bricks, concrete, metals, ceramics, plastics, sponges and plaster. Highly fluorinated polymers containing a perfluorocarbon chain linked to the polymer-chainthrough a sulfonamido group have been pre-pared by vinyl addition polymerisation of vinyl esters and allyl esters of perfluoroalkanesulphonamido - alkylenemonocarboxylic acids, as described in Patent No. 857336 and by vinyl addition polymerisation of acrylate esters and methacrylate esters of perfluoroalkanesulphonamido alkanols, as described in Patent No. 857689. However, since such 35 highly fluorinated polymers can only be dissolved in expensive fluorinated solvents at relatively low concentrations, their use in lacquers, varnishes and paints is precluded for all practical purposes.

It is, therefore, an aim of this invention to provide a fluorinated resin which can be dissolved in conventional hydrocarbon solvents to provide a clear or pigmented varnish or enamel which can be dried to an oil and water resistant coating.

The above and other aims are accomplished by the incorporation into a solvent soluble resin soluble in xylene or methyl ethyl ketone and having available hydroxyl, carboxyl, isocyanate or epoxy groups (e.g. a polyester, epoxy resin or polyurethane) of from about 1 to about 25 weight percent, based on total resin weight, of a fluorocarbon reactant having the formula:

R_f(X)_mZ or a partial ester thereof

wherein R_t is a perfluoroaliphatic (including perfluorocycloaliphatic as defined and used herein) radical having from 1 to about 18 carbon atoms, preferably 4 to 12 carbon

atoms, X-is -SO₂NR¹-or -CNR¹, m is 0 60

or 1, Z is a -R2CH2OH, or -R2COH and is connected to the nitrogen valency, wherein R2 is a substituted or unsubstituted aliphatic radical having from 1 to 18 carbon atoms, usually from 1 to 12 carbon atoms, and Ri is hydrogen or an alkyl radical having from 1 to 12, preferably from 1 to 6, carbon atoms. Based on the incorporation of the perfluoroaliphatic radical into the resin system, the total elemental fluorine content introduced into the final resin is between about 0.5 and

about 20 weight percent of the total resin solids. Significantly higher elemental fluorine content tends to adversely affect the air drying properties and the solubility of the modified resin products in the conventional hydrocarbon solvents, thereby decreasing their usefulness in varnishes, lacquers, paints and other solvent based coating compositions. However, it is also possible to dissolve some of the modified resin products in selected other solvents and also to disperse or, with the more water soluble products, to solubilise them in water.

These fluorocarbon reactants are chemically combined with and thus are incorporated into the above resins. This can be accomplished through the condensation of their hydroxyl or carboxyl groups with available carboxyl and hydroxyl groups in the resin system to form an ester linkage or bridging radical, or through the addition of their hydroxyl or carboxyl groups to an oxirane ring (resulting in an ether or ester linkage), or through the addition of their hydroxyl or carboxyl groups to an isocyanate group to form a urethane linkage and amide linkage respectively. The reaction of these fluoro-carbon acids and alcohols in such resin systems is carried out in a manner similar to that conventionally employed with nonfluorinated carboxyl or hydroxyl containing components, preferably at a temperature above 130° C.

Monofunctional alcohols useful in this invention include the N-alkanol perfluoro-alkane-sulphonamides described in U.S. Patent No. 2,803,656, which have the general formula:

R_tSO₂N(R¹)R²CH₂OH

wherein R, is a perfluoroalkyl group (including perfluorocyclcalkyl) having 4 to 12 carbon atoms, R2 is an alkylene radical having 1 to 12 carbon atoms and R1 is a hydrogen atom or an alkyl group containing 1 to 6 carbon atoms. These monofunctional alcohols are prepared by reactions of an acetate ester of a halohydrin with a sodium or potassium salt of the corresponding perfluoroalkanesulphonamide. Illustrative alcohols include the following:

N-ethyl N-ethanol perfluorooctanesulphon-

N-hexyl N-ethanol perfluorooctanesulphonamide,

N-propyl N-ethanol perfluorooctanesulphonamide,

N-ethyl N-ethanol perfluoroethanesulphon-

N-ethyl N-ethanol perfluorododecanesulphonamide,

N-hexadecyi N-ethanol perfluorooctanesulphonamide,

N - ethyl N - ethanol perfluorocyclohexylethanesulphonamide,

N - propyl N - ethancl perfluorobutylcyclohexanesulphonamide,
N - ethyl N - ethanol perfluoro-4-dodecyl-

cyclohexanesulphonamide,

- ethyl N - ethanol perfluoro - 2 methylcyclohexanesulphonamide,

N - ethyl N - hexanol perfluorooctanesulphonamide,

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N - methyl N - undecancl perfluorooctanesulphonamide,

N - methyl N - butanol perfluorobutanesulphonamide,

N - ethanol perfluorooctanesulphonamide,

Other useful monofunctional alcohols include the N-alkanol perfluoroalkanecarbexamides of the formula

R₁CN(R¹)R²CH₂OH

as described in U.S. Patent No. 2,764,602, e.g. N-ethanol perfluoroheptanecarboxamide, which may be prepared by reacting the alkanolamine with the corresponding perfluoroalkanecarboxylic acid ester, refluxing in

solvent and recovering the product.
Still other useful alcohols include the perfluoroalkyl-substituted alkanols of the formula 90

$C_nF_{2n+1}CH_2OH$,

where n is 3 to 12, (e.g. C₂F₇CH₂OH), described in U.S. Patent No. 2,666,797, and of the formula

$$R_{1}$$
— $(CH_{2})_{m}$ — OH

where R_t is a perfluoroalkyl radical having from 3 to 12 carbon atoms and m is an integer from 3 to 12

(e.g. C₈F₁₇CH₂CH₂CH₂OH,

C₃F₇CH₂CH₂CH₂OH,

C₈F₁,CH₂CH₂CH₂CH₂CH₂OH, etc.)

described in Patent No. 904263. The perfluoroalkyl-substituted alkenols may also be employed, i.e. $C_nF_{2n+1}(C_mH_{2m-2})OH$ where n is 1 to 18 and m is 1 to 18, preferably 1 105 to 12 e.g. C₈F₁₇CH=CHCH₂OH, which are described in Patent No. 904263. Further useful monofunctional alcohols include the N - (polyoxa - alkyl) - perfluoroalkane sulphonamides of Patent No. 904111, such as 110

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 $C_8F_{17}SO_2NH(CH_2)_3$ — $(OCH_2CH_2)_5$ — OH_7 C_8F_1 , $SO_2N--C_2H_4$ —(OCH₂CH)₁₀OH, etc. Ċ₂H" ĆH₃

The carboxyl-containing fluorocarbon reactants include the monofunctional perfluoroalkanesulphonamidoalkylenecarboxylic of Patent No. 857335, which have the general formula:

R₁SO₂N(R¹)R³COH

wherein R_t is a perfluoroalkyl (including perfluorocycloalkyl) groups having from 4 to 12 carbon atoms, R1 is hydrogen or an alkyl group having from 1 to 6 carbon atoms and R³ is an alkylene group having from 1 to 18 carbon atoms. These acids may be obtained by preparing a perfluoroalkanesulphonamide, converting this to a sulphonamide salt, converting the latter to an ester of the desired acid and hydrolysing the ester to form the. acid. Illustrative acids include the following:

N - cthyl N - perfluorooctanesulphonyl.

N - perfluorooctanesulphonyl glycine, N - perfluoropentanesulphonyl glycine, N - perfluorodecanesulphonyl glycine, 3 - (perfluorooctanesulphonamido) pionic acid,

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11 - (N - methyl N - perfluorooctanesulphonamido) hendecanoic acid, 18 - (N - methyl N - perfluorooctane-

sulphonamido)stearic acid, 11 - (N - ethyl N-perfluorooctanesulphon-

amido) hendecanoic acid, N - ethyl N - perfluorocyclchexylsul-

phonyl glycine, N - ethyl N - perfluorocyclohexylethane-

sulphonyl glycine,
N - butyl N - perfluoro - 4 - dodecylcyclohexanesulphonyl glycine.

N - ethyl N - perfluoro - 2 - methylcyclohexanesulphonyl-glycine,

N - hexyl N - perfluorooctanesulphenyl

glycine, N - ethyl N - perfluorobutanesulphonyl glycine.

Other carboxyl containing fluorocarbon reactants include the monofunctional perfluoroalkanecarboxamido-alkylenecarboxylic acids.

C,F₁₃CNH(CH₂)₃COOH,

which may be prepared as follows:

To a solution of 14.3 grammes (0.11 moles) of ethyl y-aminobutyrate in 100 cc of ether is added 43.2 grammes (0.1 moles) of

C₇F₁₅COOC₂H₅.

After the mixture is allowed to stand overnight, the solvent is removed under reduced pressure. The resulting ethyl ester is hydrolysed by warming in dilute sedium hydroxide solution, and the acid product is precipitated by acidification with dilute hydrochloric acid. Still other carboxyl containing fluorocarbon

reactants include the perfluoro-substituted aliphatic acids, described in Patent No. 896015, such as

C₈F₁₇CH₂CH₂CH₂CH₂COOH,

15 - perfluorobutyl pentanoic acid, 11 - perfluorooctyl-hendecanoic acid, etc. as well as the unsaturated perfluoroalkane aliphatic acids, e.g. R₁CH=CH—(CH₂)₇CH₂CO₂H, also described in U.S. Patent No. 2,951,051.

These fluorocarbon reactants may be introduced during the resin preparation or may be reacted with available oxirane rings or with free hydroxyl carboxyl or isocyanate sites on the partially or fully polymerised resin. When the proportions of the respective reactants are selected to provide an excess of available hydroxyl groups, the above fluorocarbon reactants having carboxyl substituted 80 groups are preferably employed. However, the hydrcxyl containing fluorocarbon alcohols may be utilised in such circumstances if they are reacted with dicarboxylic acids in such ratio as to form the corresponding half-ester, then the resulting half-ester is condensed with the available hydroxyl groups in the system. The former procedure is preferred, since it results in a minimum of ester linkages which may tend to hydrolyse upon extended exposure to moisture. When an excess of isc-cyanate or carboxyl groups exists in the system, the hydroxyl containing or alcoholic fluorocarbon reactants of the abovementioned type are preferably employed, although the carboxyl containing fluorocarbons-may-be-used if the partial ester with a polyhydroxy compound, e.g. trimethylol propane, ethylene glycol, etc. is first formed

In general, the resins which are modified 100 in accordance with this invention must be solvent soluble, i.e. they must be soluble in either xylene or methylethyl ketone at 70° F. to the extent of 25 weight percent or more when in unmodified form. It is therefore 105 preferred to maintain the overall average molecular weight of the unmodified resins below a maximum of about 5,000, usually below 3,500. These resins are prepared from a reactive system which contain compounds capable of either condensation polymerisation

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or non-vinyl addition polymerisation inter se and which further contain available groups reactive with active hydrogen in excess of that required for said condensation or non-

vinyl addition polymerisation.

Polyester resins having available hydroxyl or carboxyl groups may be modified with the fluorocarbon reactants of this invention. Polyester resins, as used herein, include the alkyd 10 resins, and the reactants may be either saturated or unsaturated. Since linear polyesters are produced by reacting a dibasic acid and a glycol, the sole reactive sites for chemical incorporation of the fluorocarbon reactants are at the ends of the linear chain. In such cases, the monofunctional fluorocarbon reactants serve as chain terminating agents when included in the reactant mixture, thereby limiting the molecular weight of the resinous product. If the fluorocarbon reactant is added after or during the later stages of the polymerisation, the higher molecular weights can be realised, though the maximum percent of fluorine in the modified resin is accordingly reduced. It is most preferable to include a polyfunctional acid or alcohol having at least three acidic or alcoholic groups, e.g. glycerol, pentaerythritol, trimesic acid, tricarballylic acid, etc. among the reactants to provide reactive sites along the linear chain, both for purposes of crosslinking and for esterification with the fluorocarbon acids or alcohols. Phthalic anhydride, for example, may be used with the higher 35 functional alcohols such as glycerol to produce resins having a crosslinked structure. By using a mixture of alcohols or acids, the resin properties may be usefully modified. For example, if some acid (or anhydride) such as phthalic anhydride is replaced with a relatively long chain fatty acid, such as lauric or linoleic acid, the distance between reactive or crosslinkable sites is greater and a softer resin results. If the more highly soluble glycols are used, the resin may have varying degrees of water solubility or dispersibility.

The alkyd resins are polyesters which are frequently oil modified with drying, semidrying and non-drying oils to improve flexibility. Air drying alkyds include such good drying oils as linseed and dehydrated castor oil, and semi-drying soybean oil is useful in medium and short oil alkyds. Moreover, poly-55 esters containing unsaturated acids, such as maleic or fumaric, may be modified by reacting unsaturated compounds such as styrene, methyl methacrylate, fluorinated acrylates and methacrylates (e.g. those set forth in Patent No. 857689) or fluorinated vinyl esters (e.g. those described in Patent No. 857336), with the polyester unsaturation to increase the hardness of the resin. The method for the preparation of such polyesters in both modified and unmodified form is well known and need not be further elaborated here. Examples 3, 4, 9 and 10 herein illustrate polyesters modified with the fluorocarbon reactants in accordance with this invention.

The epoxy resins, as typified by the diglycidyl ether of bisphenol A and higher homologues thereof, have a chain structure which contains ether oxygen, reactive hydroxyl groups and may also contain epoxy groups. Such resins are frequently modified with anhydrides or acids, particularly vegetable and monobasic fatty acids, by reaction with the hydroxyl groups or the epoxy groups, to provide epoxy ester resins suitable for a particular use, e.g. in air drying coatings. By employing the above described fluorocarbon acids, or a 1:1 reaction product of the above described fluorocarbon alcohols with a dibasic acid (e.g. phthalic anhydride) fluorinated epoxy esters may be prepared which have good oil and water resistance without sacrificing desirable film properties. Also, the reaction of the fluorocarbon alcohols directly with the epoxy groups proceeds rapidly in the presence of catalysts, as with the similar reaction using ordinary unfluor-inated alcohols. In one preferred illustrative embodiment a liquid epoxy resin (the diglycidyl ether of bisphenol A, epoxy equivalent 170—190) may be reacted under basic conditions with stoichiometric amounts of bisphenol A or a saturated or unsaturated dicarboxylic acid (e.g. diphenic, adipic, maleic or phthalic), a small quantity of vegetable acid to control molecular size and the desired amount of fluorocarbon acid, then the residual hydroxyl groups may be esterified with further vegetable acids (e.g. linseed oil acid) until a low acid number is attained. Solid epoxy resins which may be modified with the fluorocarbon acids or alcohols adducts may range from about 500 to about 5000 in molecular weight and have epoxide equivalents from about 350—2500. When the esterified epoxy resin contains less than about 25 percent combined weight of the fluorocarbon reactants of this invention and further is partially esterified with unsaturated acid, the modified resin has air drying properties similar to the air drying alkyds. Larger amounts of the fluorocarbon reactants tend to decrease air drying properties and adversely affect solubility of the resins in the conventional hydrocarbon solvents.

Other resins may also be modified with 120 the abovementioned fluorocarbon reactants. These resins are formed as reaction products of a diisocyanate, such as toluene diisocyanate, with an organic compound having at least two active hydrogen atoms per molecule, "active hydrogen" being defined by Zerewitinoff in Ber. 40,2025 (1906) JACS 49, 2818 (1927) and referring usually to the hydrogen cf —OH, —COOH and —NH₂ groups. Both linear and crosslinked poly- 130

urethanes may be prepared, the latter being provided by incorporation of a trifunctional reactant, such as a polyether triol (e.g. the condensation product of propylene oxide and 1,2,6-hexanetricl), into the resin system. The reactants may be varied to alter the resin properties, and the fluorocarbon reactant is incorporated into the system accordingly. Thus, in the presence of an excess of the isocyanate radical the monofunctional fluorocarbon reactants of this invention, which contain active hydrogen, add to the available isocyanate groups in the system. The fluorocarbon alcohols react to form a urethane 15 linkage, and the fluorocarbon acids react to form an amide linkage with the liberation of carbon dioxide. These fluorocarbon reactants may be first reacted with diisocyanate to form the 1:1 adduct, which can then in turn be 20 reacted with available hydroxyl groups in the resin system, as an alternative means for incorporation. If the fluorocarbon acid derivative is used in excess of the amount required for the addition reaction, a direct esterification with available hydroxyl groups (for example, as found on a polyether triol) can also be effected, although the presence of ester linkages is not generally preferred because of their tendency to hydrolyse. Hydroxyl-containing oils such as castor oil (pigmented or clear) may be included in the resinous system to add, through their hydroxyl groups, to available isocyanate groups or, if desired, such oils may be added to the moisture free system before use to lengthen shelf life. These systems may be air cured at room temperature or may be more rapidly cured at elevated temperatures in the presence of moisture.

The following examples will serve to illustrate the invention but, particularly in view of the many obvious modifications thereof, are not intended to be limiting.

Example 1

This example illustrates the fluorecarbon. modified alkyd resins of an oxidising type and a technique for their preparation. The stepwise procedure is essentially a typical alkyd cook, using the azeotropic distillation of the water in the well-known solvent

A 2-litre Pyrex glass flask equipped with thermometer, stirrer and reflux-distilling head was charged with 226 grammes of phthalic anhydride, 95 grammes of N-ethyl perfluoro-octanesulphonamido ethanol, 238 grammes of dehydrated castor oil acids and 35 grammes of xylene. The mixture was heated with stirring to 160° C. and held at this tem-.60 perature for one hour to insure essentially complete reaction of the fluorocarbon alcohol. Then 129 grammes of glycerol (a 10% excess) was added and the temperature increased to 190° C. The xylene and water of esterifica-

tion refluxed and collected, and the temperature rose to 210° C. The total processing required 51 hours to attain an acid number of 10 based on resin solids. After thinning the resinous product with xylene to 50% solids, metallic driers (cobalt and lead naphthenate) were added in concentration of 0.03% cobalt metal and 0.40% lead metal based on the weight of oil solids to aid in exidation of the eil constituent. About 10% by weight of solids of butylated melamineformaldehyde resin may be added to increase surface hardness, mar resistance and flow properties. The resultant varnish could be applied to a glass or metal surface by spraying or dip coating or by a draw down bar and curing thereon by heating in air at 300° F. for 15-45 minutes. The varnish was readily pigmentable. Flucrinated alcohol in this resin constituted 15% of resin solids and the C15 oxidising oil acid constituted 40% by weight of resin solids.

Other modified alkyd resins were prepared with variations in oil length as well as type and amount of fluorocarbon reactant. As with ordinary alkyd resins the amount and type of solvent used to thin the resin is dependent on the amount of oil acid contained in the resin. For example, aliphatic solvent such as mineral spirits and heptane are used in higher amounts of oil, and aromatic solvents such as xylene and toluene are used with lower oil content.

Example 2

This example described the preparation of a fluorocarbon modified alkyd resin containing occount oil acids (40% by weight of resin solids) a non-oxidising type because of the low degree of unsaturation of coconut oil acids. Such alkyds are usually formulated to react to a relatively high acid number (50 to 100) and are then further crosslinked with a reactive resin, e.g. butylated melamineformaldehyde, with heat curing. Non-oxidising alkyds are noted for their lack of discolouration on aging and high gloss retention 110 and are extremely useful in white baking enamels. The fluorocarbon modified nonoxidising alkyds may be prepared by the procedure of Example 1, in which the fluorocarbon alcohol is reacted with phthalic anhydride before addition of the glycerol. However, the following alternative procedure involves preparation of the alkyd resin (acid number of about 100 or higher) and subsequent refluxing with the fluorocarbon alcohol until the decreased acid number indicates essentially complete esterification of the fluorocarbon reactant.

A 1-litre Pyrex glass flask was charged with 250 grammes of an alkyd resin (non-oxdising oil, acid number 120, 80% solids in Cellosolve acetate; Cellosolve is a Reg. Trade Mark) and 66 grammes of N-ethyl perfluoro-

octanesulphonamido ethanol. The clear solution was refluxed for 2½ hours at 200° C. until the acid number was 48. Forty grammes of xylene and 164 grammes of a 55% solids solution of butylated melamine-formaldehyde resin were added. The final solids content was 70%. Hard, smooth films prepared by curing for 45 minutes at 300° F. displayed good oil and chemical resistance.

In the above alkyd examples various polyols may be substituted for glycerol, including pentaerythritol and other trihydric, tetrahydric or other polyhydric alcohols. corresponding flucrocarbon acids may be substituted for the fluorocarbon alcohols if the alkyd resin has available esterifiable hydroxyl

groups.

Example 3

A reactive polyester system was modified with the fluorocarbon reactants of this invention as follows:

A 1-litre Pyrex glass flask equipped with stirrer, thermometer and reflux distillation head was charged with 222 grammes of phthalic anhydride, 147 grammes of malcic anhydride, 114 grammes of propylene glycol, 65 grammes of N-propyl perfluoroctane-sulphonamido ethanol, and 10 grammes of toluene. After heating to 205° C. and refluxing for 2½ hours, twenty grammes of water collected. The reaction mass was then cooled to 170° C. and an additional 114 grammes of propylene glycol was introduced into the system. The temperature was increased to 210° C. and refluxing continued for 2½ hours until the acid number was 37. When the polyester had cooled to room temperature, sufficient inhibitor free styrene was added to provide a 1:1 ratio by weight of polyester/ styrene. Approximately, one gramme of benzoyl percxide was dissolved in 99 grammes of the reactive system, and films were prepared with a draw down bar. After curing for one-half hour at 140° C. the film dis-45 played excellent resistance to both water and oil. The fluorocarbon reactant constituted 5 weight percent of the polyester-styrene system.

The reactive polyester (60 grammes) was also dissolved in 40 grammes of methyl methacrylate and similarly copclymerised in the presence of a free radical catalyst or initiator (1% benzoyl peroxide). The resultant cured films were similar in water and oil resistant properties to the above polyester-styrene copolymer films.

In both of the above cases, the presence of the fluorochemical enhanced the reaction of the styrene and methacrylate with the polyester by excluding air and reducing the evaporation rate of the styrene and methacrylate monomers from the flowed out film.

Example 4

The flucrocarbon modified alkyd of Example 1 can be styrenated to improve such

properties as surface hardness of film and air 65 curing rate.

A 1-litre Pyrex glass flask equipped with dropping funnel, thermometer, stirrer and reflex condenser was charged with 320 grammes of a 75% solution of the Example 1 alkyd resin (40% oil, 15% fluorocarbon) in xylene. After warming to 90° C., 104 grammes of styrene and 2 grammes of benzoyl peroxide was added slewly over a 15-minute period. The temperature was then reject to period. The temperature was then raised to 140° C. and held for one hour. Cobalt naphthenate (0.03% cobalt based on oil weight) was added during the down heat. The final product had a clear, light brown appearance and displayed excellent shelf life. Oil and water resistant properties were excellent.

Sixty grammes of the alkyd resin was dissolved in and copolymerised with methyl

methacrylate with similar results.

Example 5

This example illustrates a typical epoxy pelyester prepared from an epoxy resin having an equivalent weight of about 150, i.e. 280 grammes of C18 fatty acid (1.0 equivalent) is required to esterify 150 grammes of epoxy resin. The fluorocarbon reactant, Nethyl perfluorooctanesulphonyl glycine, provided a 5% fluorocarbon content based on the tetal resin solids. The drying oil constituent, dehydrated castor acids, constituted

43% of the resin solids. A 1: litre Pyrex glass flask equipped with stirrer, thermometer and inert gas sparge was charged with 150 grammes of Ep 201 (3,4epoxy - 6 - methylcyclohexylmethyl - 3,4 - 100 epoxy - 6 - methylcyclohexane carboxylate; 150 equivalent weight; a product of Union Carbide Chemical Co.) 128 grammes of dehydrated castor acids and 15 grammes of N-ethyl perfluorooctanesulphonyl glycine. The 105 mixture was heated to 180° C. with stirring and the reaction was followed by acid number determinations. After one hour the acid number was 10. The product was allowed to cool to room temperature, and 33 grammes of xylene was added to make 90% solids. A catalyst sclution of BFa - etherate in xylene (4.5 grammes BF, in 213 grammes xylene) was added slowly over a one-half hour period. The solids concentration was 55% at this 115 point. A cool water bath was required to maintain a temperature range of between 25-30° C. over a 1 hour period with stirring. The addition of BF_a effects the polymerization of residual unreacted epoxy and hydroxyl groups. Water (1% based on solids) was then added to hydrolyze the boron-polymer complexes which are also formed during the polymerization. Additional xylene was added to reduce the solids content to 50%. The somewhat muddy product was treated with

sodium carbonate (2% of solids) to improve

the colour and was then filtered. Cobalt

naphthenate (0.03% cobalt metal based on weight of oil) was introduced, and films were prepared on glass and metal with a draw down bar. The resulting films heat and air cured to hard water and oil resistant films. Ten percent of a melamine-formaldehyde resin may be added to improve the surface hardness of the cured film.

Example 6

10 Using the same procedure and apparatus as in Example 5 the following reactants were employed: 245 grams of epoxy resin (eq. wt. 150), 180 grams of soya bean oil acids and 75 grams of N-ethyl perflucrooctanesulfonyl glycine. The final product contained 15% of the fluorocarbon reactant based on total solids, and was somewhat dark in appearance.

EXAMPLE 7

The dark colour of the product in Example 6 was attributed to the fluorocarbon glycinc acid. This example illustrates the use of a fluorocarbon alcohol to achieve improved pro-

duct properties.

A 1-litre Pyrex glass flask equipped with stirrer, reflux condenser and thermometer was charged with 50 grams of succinic anhydride, 288 grams of N-ethyl perfluorooctanesulfonamido ethanol and 144 grams of toluene. The solution, having an initial acid number of 130, was refluxed at 115° C. For 3½ hours, and the toluene was then removed by vacuum distillation. The product, mostly the half ester of succinic acid, had an acid number of 90. Heating was continued with stirring for another 12 hours with virtually no solvent until the final acid number was 78. The product was a white, hard waxy solid with an equivalent weight of 715 based on acid number.

Using the apparatus and procedure of Example 5 215 grams of epoxy resin (e.g. eq. wt. 150), 146 grams of soya bean oil acid, and 83 grams of the above half-ester succinic derivative were charged and reacted. The final product (55% solids in xylene) had a lighter colour, was faster drying and produced harder films than the product of the preceding example. The fluorocarbon reactant con-

stituted 15% of the final solids.

EXAMPLE 8

This example illustrates the modified polyurethane resins of this invention.

A dry 1-letre glass flask equipped with reflex condenser, stirrer, dry nitrogen gas inlet and thermometer was charged with 32.4 grams of N-ethyl perfluorooctanesulfonamido ethanol and 70.4 grams of Niax Triol LK 380, a trihydroxyl polyether (M.W. 440) prepared by reacting propylene oxide and 1, 2, 6-hexane triol (a product of Union Carbide Chemical Co.). The fluorinated alcohol dissolved in the trihydroxy polyether at 40° C.

With the temperature held at 60° C., 76.2 grams of toluene diisocyanate was introduced slowly over ½ hours into the mixture through a dropping funnel equipped with a drying tube. The temperature of the reaction mixture was increased to 80° C, with a hot water bath. To decrease the viscosity 66 grams of water-free ethyl acetate (urethane grade) was added to the flask. After maintaining the temperature, first at 80° C. for 3 hours than at 90° C. for 1½ hours, the product was thinned to 70% solids in ethyl acetate and discharged into a thin can covered with a layer of dry nitrogen. The combined fluorocarbon alcohol constituted 16% by weight of solids and the ratio of isocyanate to hydroxy groups was 2.0.

Castor oil (79 grams) was mixed into this product (358 grams) to yield a final product with 50% solids and with a ratio of isocyanate to hydroxyl of 1.5. This product can be further cured at room temperature with atmospheric moisture at 175° C. for 30 minutes. This product contained 12% by

weight of the fluorocarbon alcohol.

If a pigmented enamel is desired, the castor oil may be pigmented with a conventional enamel type pigment and thinned with water-free toluene or xylene to obtain the appropriate viscosity. The final product had good shelf life and provided a film with good oil and water repellency; particularly if the ratio of isocyanate to hydroxyl groups was in the 1.0—1.9 range. The fluorecarbon alcohol content was from about 5 to 25 percent by weight of total resin solids. Example 9

To a 1-litre, three-necked flask, equipped 100

with stirrer, thermometer, nitrogen gas inlet, reflux condenser and dropping funnel was charged 250 grams of the fluorinated alkyd of Example 1 in 50% solids solution in xylene, and 5 grams of a 73% solution of cumene hydroperoxide. The alkyd/peroxide solution was heated gradually to 80° C. with solution was neared gradually to 80°C. What stirring and with a slow flow of nitrogen. 166 grams of a 50% solution (in methyl isobutyl ketone) of N-ethyl perfluorooctane-110-sulfonamidoethyl methacrylate was slowly added at a rate of 2—3 millilitres per minute, and the solution temperature was slowly increased to reflux (130° C.) during the one hour addition period. After the addition was com- 115 plete, the reflux temperature was maintained until the viscosity increase stopped (about 6—7 hours). The light coloured resin was allowed to cool, then was placed in storage. Cobalt naphthenate (6% solution at 0.03% cobalt ion concentration was added during the down heat). The final overall fluorocarbon content was about 55% (including 15% fluoroinated alcohol reactant in fluorinated alkyd of Ex. 1). Films cured at 350° F.

resistance and were exceptionally smooth.

for 45 minutes were hard, had excellent mar

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EXAMPLE 10

Twenty grams of N-ethyl perfluorooctanesulfonamidoethyl methacrylate was dissolved in 120 grams of a 50% solution (xylene) of a fluorinated polyester (15% fluorocarbon alcohol), prepared from a 1:1 mol ratio of maleic and phthalic anhydride esterified to an acid number of 35 with propylene glycol and N-ethyl perfluorocottancsulfonamido ethanol. Benzoyl peroxide (16 grams) was dissolved in the solution. The overall combined fluorocarbon reactants was 35%. Films were cast on glass and cured at 350° F. for one hour. The films were continuous, glossy and not quite as hard as those set forth in Example 9.

Films produced from the fluorocarbon modified resins of this invention have been evaluated in terms of film hardness, air curing rate, chemical and solvent resistance, oil and water repellency, shelf life, weatering qualities, and adhesion to various substrates. Table 1 and 11 show some of these properties in comparison to the original unmodified resins. Coatings produced with the fluorocarbon modified resins are considered to have a satisfactory balance of properties if they have a Sward hardness of 15, pass a flex-ibility test on tin panels when wrapped around one-eighth inch mandrels, have good adhesion to glass and metal (i.e. no lifting of film when scratched), display good mar resistance to scratching, exhibit good solvent resistance to aromatic and aliphatic solvents, show good chemical resistance (e.g. 5% solutions of sodium hydroxide, hydrochleric acid,

appearance. Weatherability was obtained on various fluorocarbon modified resins, the data on film deterioration (e.g. loss of gloss, checking, cracking, peeling, discolcuration, etc.) being obtained after 1600 hours exposure in the weathermeter and 3½ months outdoor exposure. The water and oil resistance of the films was obtained by measurement of con-

detergents) and have excellent general film

films was obtained by measurement of contact angles with both polar and non-polar solvents.

Results of the tests indicated that from about 1 to 25 weight percent based on total 50 resin solids, preferably from 7% to 15% of the fluorocarbon acids and alcohols herein described produces modified resins which satisfactorily meet the above standards. Such fluorocarbon modified resins have superior 55 oil, water and solvent resistance as compared to the unmodified resins. They can be readily be used in conventional lacquer, enamel and varnish formulations. Weathering data on films produced therefrom has shown that the 60 fluorocarbon content imparts no deleterious effects on film durability and in fact improves gloss retention and improves overall appearances. When the resin is cured through oxidation of an unsaturated oil, curing in air at 65 75° F. was somewhat slower with the fluorocarben modified resin, but vinyl addition type modification (e.g. styrenation, methacrylation, etc.) increased air curability considerably. Moreover, by conducting such modification with a fluorinated vinyl ester, acrylate or methacrylate, the total fluorocarbon content in the final resins system can be raised to about 50 weight percent, if desired. The fluerocarbon modified polyesters, epoxys and polyurethanes of this invention are particularly useful in varnish and enamel formulations designed for use where oil borne dirt and water resistance is required, e.g. in auto finishes and refinishes, appliances, and as a general industrial paint.

EXAMPLE 11

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An air drying alkyd resin incorporating C₈F₁₇(CH₂)₁₀COOH as the fluorocarbon meiety was prepared in the following manner. Using a 3-litre reaction flask equipped with mechanical stirrer, nitrogen inlet, thermometer and a Vigreaux colum with xylene/ water azeotrope condenser and trap, about 384 grams of phthalic anhydride, 230 grams (including 16% excess) of glycerol, 98 grams of C₅F₁₇(CH₂), COOH, 200 grams of linseed fatty acids and 70 grams of xylene were charged to the reaction vessel. The mixture was heated to reflux with good agitation under a blanket of nitrogen. Reflux started at 150° C., and water started to collect at about 170° C. Refluxing was continued for about 12 hours, after which an additional 180 grams of linseed fatty acids was slowly added. The temperature was gradually increased to 220° C. and maintained until an acid value of 15 was reached. The clear resin was thinned to 60% solids in xylol (60 gram) after the temperature had dropped to 145° C the resin had a Gardner viscosity of about L and a clear colour of about 12. A film containing the usual driers was allowed to dry overnight and was dried throughout in 2 days. The clear film had an oil centact angle of 60°.

Similar results can be achieved by replacing the saturated fluorochemical acid with

O || C,F,;C—NH(CH₂);COOH.

Example 12

About 144 grams of phthalic anhydride, 1 42 grams of glycerol, 86 grams of linseed fatty acids (alkali refined), 35 grams of

> O || C,F,,:CNHCH,:CH,:OH

and 20 grams of xylene were charged to a 1-litre 3-necked reaction flask equipped as 12 described in Example 11. The mixture was

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heated to reflux (about 140° C.) and maintained for about one hour, when an added 42 grams of glycerol was added and the temperature increased to about 170° C. During the ensuing three hours of reaction, an additional 86 grams of linseed acids was added. The maximum temperature attained was 215° C. The final acid value on solids was 25. The clear, light brown resin had a Gardner viscosity of Z as a 75% solids solution in xylene and a colour of 12. A film of the 75% solution containing conventional driers was drawn down on glass and cured one hour at 250° F., after which the film was dry but slightly tacky. The film was then cured for an additional hour at 300° F: to yield a dry, rubbery film with a hexadecane contact angle of 50° C.

Example: 13

A reaction flask equipped as in the preceding examples was charged with 35 grams of phthalic anhydride, 23 grams of glycerol, 41 grams of dehydrated castor acids and 6 grams of C₈F₁,CH=CHCH₂OH. The mixture was refluxed for 3½ hours until an acid value of 4.3 was reached. The Gardner viscosity of the 50% solids solution in xylene was E—F. A 3-mil film containing conventional driers was drawn down on glass and cured in air for 30 minutes with subsequent baking at 300° F. for one hour. A hard clear film was produced, which gave a hexadecane droplet contact angle of 50°.

Example 14

This example describes the chemical incorporation of $C_8F_{17}SO_2N(C_2H_2)CH_2CH_2OH$ in an alkyd resin that may be dispersed in water to form an aqueous system or may also be dissolved in organic solvents rather than water dispersions, if desired.

A 1-litre reaction flask fitted with a mechanical stirrer, inert gas inlet tube, thermometer and a water/xylene azeotrope head with condenser was charged with 535 grams of refined linseed acids, 185 grams of pentaery-thritol, 237 grams of phthalic anhydride, 122 grams_of_polyethylene_glycol_(average_mole_

cular weight of 1540) and 70 grams of xyl nz. The heating was effected with a heating mantle and powerstat. The mixture was heater under a blanket of nitrogen to 153° C., at which temperature refluxing started. Water of esterification also started to collect in the azeotrope separator at this point. The temperature was gradually increased over a four-hour period to a peak temperature of 243° C., collecting about 62 cc of water of esterification which was continually drawn off during the reaction. At this point, the acid value on a solids basis had fallen to 25. The solids content was about 95% in xylene. About one-half (500 g) of the alkyd solids was charged to another reaction flask of similar type. About 86 of

$C_8F_{17}SO_2N(C_2H_5)CH_2CH_2OH$

and 30 g. of xylene were charged to the flask containing the prepared alkyd. The mixture was warmed to about 100° C. to effect good mixing; the acid value on solids at this point was 21. The solution was increased in temperature to 220° C. where refluxing started. The temperature rose to 260° C. after 2 hours (peak). The temperature was decreased to 250° C. and held therefor an additional hour. At this point the acid value of the clear resin solids was 12.

When the temperature had fallen to about $90-100^{\circ}$ C., the alkyd was poured slowly into rapidly agitated hot (about 70° C.) distilled water to form a thick, milky dispersion of excellent stability. The solids content in water was $50\pm1\%$ with a Brookfield viscosity of 6000 cps. The alkyd may be cured as films with cobalt or manganese naphthenate (0.03%) metal based on oil solids). Air dry film will dry in one day and through dry in several. The alkyd resin may be blended with other types of latices and water systems. Films of the alkyd itself or in blends with other resins display oil contact angles on cured film of 70° C. or higher.

Other modifications and embodiments will be apparent from the above disclosure without departing from the scope of this invention 90

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TABLE I

Description of Resins and Physical Properties of Films

| | Denoment | | 2000 E | Contact Angle | Angle | | |
|--|----------|-----------|-------------------------------------|---------------|-------|------------------------|--|
| Resin | Fluorine | Ability | 300 F. 45 min. | *lio | Water | Application | Properties |
| Phthalic Alkyd (oxidising) | 4 | fair | Exœllent Sl. yellow | 55° | °06 | Spray, dip | Satisfactory balance |
| Phthalic Alkyd (oxidizing) | œ | Fair | excellent SI. yellow | , 65° | °06 | spray, dip | satisfactory balance |
| Phthalic Alkyd (semi-oxidizing) | 13 | poog | excellent No yellow | ,0Z | °06 | spray, dip | satisfactory balance (excellent solvent resistance) |
| Phthalic Alkyd (non-oxidizing) | 10 | i | excellent no yellow | °89 | °06 | spray, dip | satisfactory balance (excellent solvent resistance) |
| Hpoxy Polyester (oxidizing) | ∞ | fair | excellent SI. yellow | . 65°. | °06 | spray, dip | satisfactory balance |
| Phthalic Alkyd control (oxidizing) | none | pood | excellent Sl. yellow | none | °09 | spray, dip | satisfactory balance (fair solvent resistance) |
| Polyurethane | 9 | excellent | excellent (200° F.) No yellow | 45° | . 66° | spray, dip | satisfactory balance (excellent solvent resistance) |
| Polyurethane | none | excellent | excellent | none | 09 | spray, dip | satisfactory balance |
| Alkyd, Styrenated | 4 | very good | excellent SI. yellow | .02 | °06 | spray, dip | satisfactory balance |
| Polyester — styrene 1% MEK*** peroxide | 4 | fair | pood | 20° | °06 | spray, curtain coat | satisfactory balance |

* Hexadecane

^{**} Percent Fluorine (elemental) is approximatedly one-half of the weight of the fluorocarbon derivative

^{***} MEK = methyl ethyl ketone

TABLE II

Weatherability (Weatherometer and Outdoor)*

| Resin | Percent Fluorine | Exposure | Checking/ Peeling | Fading | Gloss | General Appearance |
|-------------------|---------------------|---------------|----------------------|------------|-------|-----------------------|
| Alkyd (oxidizing) | 4 | none | 5 | 2 | . 5 | 20 |
| Alkyd (oxidizing) | 4 | weatherometer | ĭ | 4 | ю | ₹ |
| Alkyd (oxidizing) | 4 | outdoor | ۍ | 4 | ` m | m |
| Alkyd (oxidizing) | none | none | | 1 0 | 5 | īC |
| Alkyd (oxidizing) | none | weatherometer | ۲. | 4 | m | m |
| Alkyd (oxidizing) | none | outdoors | س | 4 | . 23 | <u>ښ</u> |

Above panels were pigmented with titanium dioxide, phthalocyanine blue or toluidine red in percentages of 20% PVC (pigment volume concentration)

* Weatherometer 1000 hours, outdoor St. Paul, Minn 45° to the South

5 — excellent

4 — good

3 — fair

2 — poor

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WHAT WE CLAIM IS:-

1. A modified resin comprising a resin soluble in either xylene or methyl ethyl ketone and having bonded thereto from about 1 to about 25 weight percent, based on total resin weight, of a fluorochemical substituent having the formula

R₁(X)_mZ or a partial ester thereof

wherein R, is a perfluorinated aliphatic 10 radical having from 1 to about 18 carbon atoms, R₂X is selected from the group consisting of

m is 0 or 1, R¹ is selected from the group consisting of hydrogen and an alkyl radical having from 1 to 12 carbon atoms, and Z is selected from the group consisting of R²CH₂OH and R²COOH, where R² is selected from the group consisting of substituted or unsubstituted aliphatic radicals having from 1 to 18 carbon atoms, said bonding being effected through a bridging radical selected from the group consisting of

the elemental fluorine content of said modified resin attributed to the R_t radical being from 0.5 to about 20 weight percent of total

resin.

2. A modified resin which comprises a resin selected from the group consisting of a condensation polyester resin, an epoxy resin and a polyurethane resin having bonded thereto from about 1 to about 25 weight percent, based on total resin weight, of a fluorochemical substituent having the formula

R_f(X)_mZ or a partial ester thereof

wherein R₁ is a perfluoroaliphatic radical having from 1 to about 18 carbon atoms, R₂X to is selected from the group consisting of

m is 0 or 1, R¹ is selected from the group consisting of hydrogen and an alkyl radical having from 1 to 12 carbon atoms, and Z is

selected from the group consisting of

R°CH2OH and R°COOH,

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where R² is selected from the group consisting of substituted or unsubstituted aliphatic radicals having from 1 to 18 carbon atoms, said bonding being effected through a bridging radical selected from the group consisting

the elemental fluorine content of said modified resin attributed to the R₁ radical being from 0.5 to about 20 weight percent of total resin

3. A modified resin suitable for use in paint, varnish lacquer and the like which comprises a condensation polyester resin having bonded thereto from about 1 to about 25 weight percent, based on total resin weight, of a fluorocarbon having the formula

$$R_i(X)_mZ$$
 or a partial ester thereof

wherein R_t is a perfluoroaliphatic radical having from 1 to about 18 carbon atoms, R_tX is selected from the group consisting of

m is 0 or 1, R¹ is selected from the group consisting of hydrogen and an alkyl radical having from 1 to 12 carbon atoms, and Z is selected from the group consisting of

R2CH2OH and R2COOH,

where R² is selected from the group consisting of substituted or unsubstituted aliphatic radicals having from 1 to 18 carbon atoms, said bonding being effected through ester linkages.

4. A modified resin suitable for use in paint, varnish lacquer and the like which comprises an epoxy resin having bonded thereto from about 1 to about 25 weight percent, based on total resin weight, of a fluorocarbon having the formula

R_f(X)_mX or a partial ester thereof

wherein R_r is a perfluorinated aliphatic radical having from 1 to about 18 carbon atoms, $R_t X$ is selected from the group consisting of

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O || |R_iSO₂NR¹ and R_iCNR¹

m is 0 or 1, R¹ is selected from the group consisting of hydrogen and an alkyl radical having from 1 to 12 carbon atoms, and Z is selected from the group consisting of

R2CH2OH and R2COOH,

where R² is selected from the group consisting of substituted or unsubstituted aliphatic radicals having from 1 to 18 carbon atoms, said bonding being effected through ester linkages.

5. A modified resin suitable for use in paint, varnish lacquer and the like which comprises a polyurethane resin having bonded thereto from about 1 to about 25 weight percent, based on total resin weight, of a fluorocarbon having the formula

R₁(X)_mZ or a partial ester thereof

wherein R_t is a perfluorinated aliphatic 20 radical having from 1 to about 18 carbon atoms, R_tX is selected from the group consisting of

Q || |R₂SO₂NR¹ and R₂CNR¹

m is 0 or 1, R¹ is selected from the group consisting of hydrogen and an alkyl radical having from 1 to 12 carbon atoms, and Z is selected from the group consisting of

R2CH2OH and R2COOH,

where R² is selected from the group consist-30 ing of substituted or unsubstituted aliphatic radicals having from 1 to 18 carbon atoms, said bonding being effected through a linkage selected from the group consisting of a urethane, ester, and amide group.

6. A polymerizable system comprising reactive compounds being capable of condensation polymerization inter se and having available groups reactive with active hydrogen in excess of that required for said condensation polymerization, and, as a modifying ingredient, between about 1 and about 25 weight percent, based on said reactive compounds, of a fluorochemical compound having the formula

45 R₁(X)_mZ or a partial ester thereof

wherein R_t is a perfluorinated aliphatic radical having from 1 to about 18 carbon

atoms, R_iX is selected from the group consisting of

R₁SO₂NR¹ and R₁CNR

m is 0 or 1, R¹ is selected from the group consisting of hydrogen and an alkyl radical having from 1 to 12 carbon atoms, and Z is selected from the group consisting of

R2CH2OH and R2COOH,

where R² is selected from the group consisting of substituted or unsubstituted aliphatic radicals having from 1 to 18 carbon atoms.

7. A polymerizable system comprising reactive compounds being capable of non-vinyl addition polymerization inter se and having available groups reactive with active hydrogen in excess of that required for said non-vinyl addition polymerization, and, as a modifying ingredient, between about 1 and about 25 weight percent, based on said reactive compounds, of a fluorochemical compound having the formula

R_f(X)_mZ or a partial ester thereof

wherein R_t is a perfluorinated aliphatic radical having from 1 to about 18 carbon atoms, X is selected from the group consisting of

O || R₁SO₂NR¹ and R₁CNR

m is 0 or 1, R¹ is selected from the group consisting of hydrogen and an alkyl radical having from 1 to 12 carbon atoms, and Z is selected from the group consisting of

R2CH2OH and R2COOH.

where R² is selected from the group consisting of alkylene and alkenylene radicals having from 1 to 18 carbon atoms.

8. A process for the preparation of a fluorinated resin comprising the condensation or non-vinyl addition polymerization of reactive compounds to produce a resin soluble in xylene or methyl ethyl ketone, and the reaction therewith, either during or after polymerization at a temperature above about 130° C. of a fluorochemical compounds of the formula

R_f(X)_mZ or a partial ester thereof

wherein R_i is a perfluorinated aliphatic radical having from 1 to about 18 carbon atoms,

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R_tX is selected from the group consisting of

m is 0 or 1, R^1 is selected from the group consisting of hydrogen and an alkyl radical having from 1 to 12 carbon atoms, and Z is selected from the group consisting of

R2CH2OH and R2COOH,

where R² is selected from the group consisting of substituted or unsubstituted aliphatic radicals having from 1 to 18 carbon atoms, the reaction being effected through a bridging radical selected from the group consisting of

O and —CNH—,

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the elemental fluorine content of said modified resin attributed to the R_r radical being from 0.5 to about 20 weight percent of total resin.

9. The process of claim 8 wherein said polymerization is condensation polymerization.

10. The process of claim 8 wherein said polymerization is non-vinyl addition polymerization.

11. A modified resin substantially as herein described with reference to the Examples.

12. A precess for the preparation of a fluorinated resin substantially as herein described with reference to the Examples.

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Learnington Spa: Printed for Her Majesty's Stationery Office, by the Courier Press (Learnington) Ltd.—1965. Published by The Patent Office, 25 Southampton Buildings, London, W.C.2, from which copies may be obtained